

EXHIBIT I in USSN 09/445,517

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Related Studies

A Study to Characterize Regimens of Basal Insulin Intensified With Either Symlin® or Rapid Acting Insulin in Patients With Type 2 Diabetes

This study has been completed.

Study NCT00467649 Information provided by Amylin Pharmaceuticals, Inc. First Received: April 27, 2007 Last Updated: April 10, 2009 History of Changes

Study Type:	terventional			
Study Design:	domized, Open Label, Active Control, Parallel Assignment			
Condition:	2 Diabetes Mellitus			
Interventions:	Drug: pramlintide acetale Drug: rapid acting insufin (Humalog® (insulin lispro), Novolog® (insulin aspart), or Apidra® (insulin glulisine)) Drug: bassi insulin (Lartius® (insulin glargine), or Levernir® (insulin determir))			

Participant Flow

Recruitment Details

Key information relevant to the recruitment process for the overall study, such as dates of the recruitment period and locations

No lext enlered.

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Pre-Assignment Details

Significant events and approaches for the overall study following participant enrollment, but prior to group assignment

No text entered.

Reporting Groups	
	Description
Group A (P1 SYMLIN)	SYMLIN treatment (120 mcg prior to major meals) was initiated on Day 1. Basal insulin was titrated throughout the study
Group B (P1 RA Insulin)	Rapid acting insulin (RA Insulin: variable dosing, threted to optimize postprandial glucose control) was initiated at Week 4. Basal insulin was litrated throughout the study
Group C (P2 SYMLIN)	Patients from Group A, who achieved HbA1c goal at Week 24, continued Phase 1 treatment during Phase 2
Group D (P2 SYMLIN+RA)	Patients from Group A, who did not achieve HbA1c goal at Week 24, continued phase 1 treatment and initiated RA insufin during Phase 2
Group E (P2 RA Insulin)	Patients from Group B, who schleved HbA1c goal at Week 24, continued Phase 1 treatment during Phase 2
Group F (P2 RA Insulin + SYMLIN)	Patients from Group B, who did not achieve HbA1c goal at Week 24, continued phase 1 treatment and Initiated SYMLIN during Phase

Participant Flow for 2 periods

Period: Phase 1 (Intent-to-Treat Population)

	Group A (P1 SYMLIN)	Group B (P1 RA Insulin)	Group C (P2 SYMLIN)	Group D (P2 SYMLIN+RA)	Group E (P2 RA Insulin)	Group F (P2 RA Insulin + SYMLIN)
STARTED	56	56	0	0	0	0
COMPLETED	48	50	0	0	0	0
NOT COMPLETED	8	6	0	0	0	0
Adverse Event	2	0	0	0	0	0
Investigator Decision	1	0	0	0	0	0
Lost to Follow-up	2	4	0	0	0	0
Withdrawal of Consent	3	2	0	0	0	0

	Group A (P1 SYMLIN)	Group B (P1 RA Insulin)	Group C (P2 SYMLIN)	Group D (P2 SYMLIN+RA)	Group E (P2 RA Insulin)	Group F (P2 RA Insulin + SYMLIN)
STARTED	0	0	17	31	14	36
COMPLETED	0	0	17	29	14	35
NOT COMPLETED	0	0	0	2	0	1
Lost to Follow-up	0	0	0	1	0	0
Protocol Violation	0	0	0	1	0	0
Withdrawal of Consent	0	0	0	0	0	1

Baseline Characteristics

Reporting Groups

	Description
Group A (P1 SYMLIN)	SYMILIN treatment (120 mag prior to major masis) was initiated on Day 1. Basal insulin was fitrated throughout the study
	Rapid acting insulin (RA Insulin: vanable dosing, fitrated to optimize postprandial glucose control) was initiated at Week 4. Basal insulin was litrated throughout the study.

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Saseline Measures			
	Group A (P1 SYMLIN)	Group B (P1 RA Insulin)	Total
Number of Participants [units: participants]	56	56	112
Age [units: participants]			
<=18 years	0	0	0
Between 18 and 65 years	46	49	95
>=65 years	10	7	17
Age (units: years) Mean ± Standard Deviation	55.0 ± 11.35	53.6 ± 9.70	54.3 ± 10.53
Gender [units: participants]			
Female	22	19	41
Male	34	37	71
Region of Enrollment [units: participants]			
United States	56	56	112
Fasting Plasma Glucose [units: mg/dL] Mean ± Standard Deviation	155,1 ± 39,60	164.3 ± 49.61	159.7 ± 44.92
Fasting Serum Lipids [units: mg/dL] Mean ± Standard Deviation			
Total Cholesterol	167.53 ± 47.054	169.86 ± 49.121	168.70 ± 47.903
HDL	44.71 ± 11.893	41.77 ± 9.468	43.23 ± 10.790
LDL	89.15 ± 38.386	90.41 ± 34.114	89.78 ± 36.133
Triglycerides	174.13 ± 108.257	193.59 ± 159.508	183.95 ± 136.273
HbA1c [units: %] Mean ± Standard Deviation	8.19 ± 0.840	8.25 ± 0.816	8.22 ± 0.825
Walst Circumference (units: cm) Mean ± Standard Deviation	116.31 ± 15.427	117.15 ± 13.198	116.73 ± 14.297
Weight (units: kg) Mean ± Standard Deviation	107.87 ± 21.893	103.46 ± 17.908	105.67 ± 20.032

► Outcome Measures

Hide results for all outcome measures

1. Primary Outcome Measure: The Proportion of Patients Achieving HbA1c <=7% at Week 24 With no Gain in Body Weight From Baseline and no incidence of Severe Hypoglycemia

Measure Type	Primary	
Measure Title	easure Title The Proportion of Patients Achieving HbA1c <=7% at Week 24 With no Gain in Body Weight From Baseline and no incidence of Severe Hypoglycemia	
Measure Description	ription Comprehensive treatment endpoint assessing the achievment of glycemic control without weight gain and severe hypoglycemia. The patier must achieve each component of the endpoint to count towards the final percentage.	
Time Frame	Weeks	
Safaty legua	No.	

Population Description

Explanation of how the number of participants for analysis was determined. Includes whether analysis was per protocol, intention to treat, or another method. Also provides relevant details such as imputation technique, as appropriate.

Phase 1 Intent-to-Treat LOCF. LOCF: If a treated patient has missing result value at week 24, then last observed value before week 24 and after baseline is carried forward to impute the week 24 value.

Reporting Groups	
	Description

Group A (P1 SYMLIN)	SYMLIN treatment (120 mgg prior to major meals) was initiated on Day 1. Basal insulin was titrated throughout the study
Group B (P1 RA Insulin)	Rapid acting insulin (RA Insulin, variable dosing, litrated to optimize postprandial glucose control) was initiated at Week 4, Basal insulin was filtrated throughout the study

Measured Values

	Group A (P1 SYMLIN)	Gre B (R Insi
Number of Participants Analyzed (units: participants)	56	5
The Proportion of Patients Achieving HbA1c <=7% at Week 24 With no Gain in Body Weight From Baseline and no incidence of Severe Hypoglycemia (units: %)	30.4	10

Statistical Analysis 1 for The Proportion of Patients Achieving HbA1c <=7% at Week 24 With no Gain in Body Weight From Baseline and no Incidence of Severe Hypoglycomia

Groups [1]	All groups
Method [2]	Fisher Exact
P Value [7]	0.0180

[1] Additional details about the analysis, such as null hypothesis and power calculation: No text entered.

[2] Other relevant information, such as adjustments or degrees of freedom:

No text entered.

[3] Additional information, such as whether or not the p-value is adjusted for multiple comparisons and the a priori threshold for statistical significance: No text entered.

2. Secondary Outcome Measure: Proportion of Patients Achieving HbA1c <= 7% at Week 24

Measure Type	Secondary
Measure Title	Proportion of Patients Achieving HbA1c <=7% at Week 24
Measure Description	This is a component of the primary endpoint
Time Frame	24 Weeks
Safety Issue	No

Population Description

Explanation of how the number of participants for analysis was determined, includes whether analysis was per protocol, intention to treat, or another method. Also provides relevant details such as imputation technique, as appropriate. Phase 1 Intent-to-Treat

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	Description	
Group A (P1 SYMLIN)	SYMLIN treatment (120 mag prior to major meals) was initiated on Day 1, Basal Insulin was fitrated throughout the study	
	Rapid acting Insulin (RA Insulin: variable dosing, Strated to optimize postprandial glucose control) was initiated at Week 4. Basal insulin was strated throughout the study	

Measured Values

	Group A (P1 SYMLIN)	Group B (P1 RA Insulin)
Number of Participants Analyzed [units: participants]	56	56
Proportion of Patients Achieving HbA1c <=7% at Week 24 junits: %	44.6	55.4

No statistical analysis provided for Proportion of Patients Achieving HbA1c <=7% at Week 24

3. Secondary Outcome Measure: Proportion of Patients With no Weight Gain at Week 24

Measure Type	Secondary
Measure Title	Proportion of Patients With no Weight Gain at Week 24
Measure Description	This is a component of the primary endpoint
Time Frame	24 Weeks
Safety Issue	No

Population Description

Explanation of how the number of participants for analysis was determined. Includes whether analysis was per protocol, intention to treat, or another method. Also provides relevant details such as imputation technique, as appropriate. Phase 1 Intent-to-Treat

Reporting Groups

	Description
Group A (P1 SYMLIN)	SYMLIN treatment (120 mgg pnor to major meals) was initiated on Day 1. Basal insulin was titrated throughout the study
Group B (P1 RA Insulin)	Rapid acting insulin (RA Insulin: vanable dusing, titrated to optimize postprandlel glucose control) was initiated at Week 4. Basal insulin was titrated throughout the study

Measured Values

	Group A (P1 SYMLIN)	Group B (P1 RA Insulin)
Number of Participants Analyzed [units: participants]	56	56
Proportion of Patients With no Weight Gain at Week 24	46.4	14.3

No statistical analysis provided for Proportion of Patients With no Weight Gain at Week 24

4. Secondary Outcome Measure: Proportion of Patients With a Severe Hypoglycemia Adverse Event

Measure Type	Secondary
Measure Title	Proportion of Patients With a Severe Hypoglycemia Adverse Event
Measure Description	This is a component of the primary endpoint
Time Frame	24 Weeks
Safety Issue	No

Population Description

Explanation of how the number of participants for analysis was determined. Includes whether analysis was per protocol, intention to treat, or another method. Also provides relevant details such as imputation technique, as appropriate.

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	Description	
Group A (P1 SYMLIN) SYMLIN treatment (120 mcg prior to major meals) was initiated on Day 1. Basal insulin was titrated throughout the study		
Group B (P1 RA Insulin) Rapid acting insulin (RA Insulin: variable dosing, Bireled to optimize postprandial glucose control) was initiated at Week 4. Basal in was tritiated throughout the study		

Measured Values

ł .	Group A (P1 SYMLIN)	Group B (P1 RA Insulin)
Number of Participants Analyzed [units: participants]	56	56
Proportion of Patients With a Severe Hypoglycemia Adverse Event [units: %]	0.0	0.0

No statistical analysis provided for Proportion of Patients With a Severe Hypoglycemia Adverse Event

5. Secondary Outcome Measure: Change in HbA1c From Baseline at Week 24

Measure Type	Secondary
Measure Title	Change in HbA1c From Baseline at Week 24
Measure Description	No text entered.
Time Frame	24 Weeks
Safety Issue	No

Population Description

Explanation of how the number of participants for analysis was determined. Includes whether analysis was per protocol, intention to treat, or another method. Also provides relevant details such as imputation technique, as appropriate.

Phase 1 Intent-to-Treat LOCF. LOCF: If a treated patient has missing result value at week 24, then last observed value before week 24 and after baseline is carried forward to impute the week 24 value.

Reporting Groups

	escription	
Group A (P1 SYMUN)	SYMLIN treatment (120 mg prior to major meals) was initiated on Day 1, Basal insulin was filtrated throughout the study	
Group B (P1 RA Insulin)	Rapid acting insulin (RA Insulin: variable doxing, titrated to optimize postprandial glucose control) was initiated at Week 4. Basal insulin was titrated throughout the study	

Measured Values

	Group A (P1 SYMLIN)	Group B (P1 RA Insulin)
Number of Participants Analyzed [units: participants]	56	56
Change in HbA1c From Baseline at Week 24 [units: %] Least Squares Mean ± Standard Error	-1.11 ± 0.17	-1.27 ± 0.17

No statistical analysis provided for Change in HbA1c From Baseline at Week 24

6. Secondary Outcome Measure: Change in Body Weight From Baseline at Week 24

Measure Type	Secondary
Measure Title	Change in Body Weight From Baseline at Week 24
Measure Description	No text entered.
Time Frame	24 Weeks
Safety Issue	No

Population Description

Explanation of how the number of participants for analysis was determined. Includes whether analysis was per protocol, Intention to treat, or another method. Also provides relevant details such as imputation technique, as appropriate.

Phase 1 Intent-to-Treat LOCF, LOCF: If a treated patient has missing result value at week 24, then last observed value before week 24 and after baseline is carried forward to impute the week 24 value.

Reporting Groups

	Description
Group A (P1 SYMLIN)	SYMLIN treatment (120 mcg prior to major meals) was initiated on Day 1. Basal insulin was titrated throughout the study
Group B (P1 RA Insulin)	Rapid acting insulin (RA insulin; veriable dosing, filtrated to optimize postprandial glucose control) was initiated at Week 4. Basal insulin was titrated throughout the study

Measured Values

	Group A (P1 SYMLIN)	Group B (P1 RA Insulin)
Number of Participants Analyzed [units: participants]	56	56
Change in Body Weight From Baseline at Week 24 [units: kg] Least Squares Mean ± Standard Error	0.02 ± 0.68	4.65 ± 0.68

No statistical analysis provided for Change in Body Weight From Baseline at Week 24

7. Secondary Outcome Measure: Change in Waist Circumference From Baseline

Measure Type	Secondary	
Measure Title	Change in Waist Circumference From Baseline	
Measure Description	No text entered.	
Time Frame	24 Weeks	
Safety Issue	No	

Population Description

Explanation of how the number of participants for analysis was determined, includes whether analysis was per protocol, intention to treat, or another method. Also provides relevant details such as imputation technique, as appropriate.

Phase 1 Intentio-Treat LOCF: LOCF: If a treated patient has missing result value at week 24, then last observed value before week 24 and after baseline is carried forward to impute the week 24 value.

Reporting Groups Description

Group A (P1 SYMLIN)	SYMLIN treatment (120 mcg prior to major meals) was initiated on Day 1, Basal insulin was titrated throughout the study
Group B (P1 RA Insulin)	Rapid acting insulin (RA Insulin: variable dosing, titrated to optimize postprandial glucose control) was initiated at Week 4. Basal insulin was litrated throughout the study

Measured Values

	Group A (P1 SYMLIN)	Group B (P1 RA Insulin)
Number of Participants Analyzed [units: participants]	53	56
Change in Waist Circumference From Baseline		
(units: cm) Least Squares Mean ± Standard Error		
Change at Week 24	-0.63 ± 0.87	2.17 ± 0.86

No statistical analysis provided for Change in Walst Circumference From Baseline

8. Secondary Outcome Measure: Change in Fasting Plasma Glucose From Baseline

Measure Type	Secondary
Measure Title	Change in Fasting Plasma Glucose From Baseline
Measure Description	No text entered.
Time Frame	24 Weeks

Safety Issue	No	

Explanation of how the number of participants for analysis was determined. Includes whether analysis was per protocol, intention to treat, or another method. Also provides relevant details such as imputation technique, as appropriate.

	Reporting Groups					
Description						
Group A (P1 SYMLIN) SYMLIN treatment (120 mcg prior to major meals) was initiated on Day 1. Basal insulin was bitrated throughout the study						
Group B (P1 RA Insulin) Rapid acting insulin (RA Insulin: variable dosing, titrated to optimize postprandial glucose control) was initiated at Week 4. Bas was titrated throughout the study						

Measured Values

	Group A (P1 SYMUN)	Group B (P1 RA Insulin)
Number of Participants Analyzed [units: participants]	45	50
Change in Fasting Plasma Glucose From Baseline		
[units: mg/dL] Mean ± Standard Error		
Change at Week 24	-29.0 ± 7.32	-37.8 ± 7.69

No statistical analysis provided for Change in Fasting Plasma Glucose From Baseline

9. Secondary Outcome Measure: Fasting Serum Lipids Change From Baseline at Week 24

Measure Type	Secondary
Measure Title	Fasting Serum Lipids Change From Baseline at Week 24
Measure Description	No text entered.
Time Frame	24 Weeks
Safety Issue	No

Population Description

Explanation of how the number of participants for analysis was determined. Includes whether analysis was per protocol, intention to treat, or another method. Also provides relevant details such as imputation technique, as appropriate. Phase 1 Intent-to-Treat

porting Groups

	Description
Group A (P1 SYMLIN)	SYMLIN treatment (120 mog prior to major meals) was initiated on Day 1. Basal insulin was titrated throughout the study
Group B (P1 RA Insulin)	Rapid acting insulin (RA Insulin: variable dosing, titrated to optimize postprandial glucose control) was initiated at Week 4. Basal Insulin

	Group A (P1 SYMUN)	Group B (P1 RA Insulin)
Number of Participants Analyzed [units: participants]	47	49
Fasting Serum Lipids Change From Baseline at Week 24		
[units: mg/dt.] Mean ± Standard Error		
Total Cholesterol	-1.81 ± 5.826	5.27 ± 4.649
HDL.	1.11 ± 1.190	1.65 ± 1.075
LDL	2.36 ± 4.456	9.12 ± 3.865
Triglycerides	-28.96 ± 12.442	-31.98 ± 13.883

No statistical analysis provided for Fasting Serum Lipids Change From Baseline at Week 24

10. Secondary Outcome Measure: Phase 2: Change in HbA1c at Week 36

Measure Type	Secondary	
Measure Title	Phase 2: Change in HbA1c at Week 36	
Measure Description	No text entered.	
Time Frame	36 Weeks	
Safety Issue	No	

Population Description

Explanation of how the number of participants for analysis was determined. Includes whether analysis was per protocol, intention to treat, or another method. Also provides relevant details such as imputation technique, as appropriate.

Phase 2 Intent-to-Treat

Reporting Groups

Description

Group C (P2 SYMLN)

Patterts from Group A, who adviseed HtA1's goal at Week 24, continued Phase 1 treatment during Phase 2

Group D (P2 SYMLN-RA)

Patterts from Group A, who do not advise thA1's goal at Week 24, continued phase 1 treatment and initiated RA insulin during Phase 2

Group E (P2 RA Insulin)

Patterts from Group B, who addressed HtA1's goal at Week 24, continued Phase 1 treatment during Phase 2

Group E (P2 RA Insulin)

Patterts from Group B, who addressed HtA1's goal at Week 24, continued Phase 1 treatment during Phase 2

Measured Values

	Group C (P2 SYMLIN)	Group D (P2 SYMLIN+RA)	Group E (P2 RA Insulin)	Group F (P2 RA Insulin + SYMLIN)
Number of Participants Analyzed [units: participants]	17	30	14	36
Phase 2: Change in HbA1c at Week 36				
[units: %] Mean ± Standard Error				
Change From Baseline	-1.96 ± 0.238	-0.68 ± 0.174	-1.49 ± 0.189	-0.99 ± 0.157
Change From Week 24	0.14 ± 0.062	-0.23 ± 0.123	0.22 ± 0.097	0,07 ± 0,113

No statistical analysis provided for Phase 2: Change in HbA1c at Week 36

11. Secondary Outcome Measure: Phase 2: Change in Body Weight at Week 36

Measure Type	Secondary		
Measure Title	Phase 2: Change in Body Weight at Week 36		
Measure Description	No text entered.		
Time Frame	36 Weeks		
Safety Issue	No		

Population Description

Explanation of bow the number of participants for analysis was determined. Includes whether analysis was per protocol, intention to treat, or another method. Also provides retevant details such as imputation technique, as appropriate.

Phase 2 Intention-Treat

Reporting Groups

	Description		
Group C (P2 SYMLIN)	Patients from Group A, who achieved HbA1c goal at Week 24, continued Phase 1 treatment during Phase 2		
Group D (P2 SYMLIN+RA)	Patients from Group A, who did not achieve HbA1c goal at Week 24, continued phase 1 treatment and initiated RA insulin during Phase 2		
Group E (P2 RA Insulin)	Patients from Group B, who achieved HbA1c goal at Week 24, continued Phase 1 treatment during Phase 2		
Group F (P2 RA Insulin + SYMLIN)	Patients from Group B, who did not achieve HbA1c goal at Week 24, continued phase 1 treatment and initiated SYMLIN during Phase 2		

Measured Values

	Group C (P2 SYMLIN)	Group D (P2 SYMLIN+RA)	Group E (P2 RA Insulin)	Group F (P2 RA Insulin + SYMLIN)
Number of Participants Analyzed [units: participants]	17	30	14	36
Phase 2: Change in Body Weight at Week 36 [units: kg] Mean ± Slandard Error				
Change From Baseline	-0.80 ± 2.096	1.34 ± 0.933	3,90 ± 1,488	4.51 ± 0,761
Change From Week 24	0.69 ± 0.654	0.50 ± 0.303	0.44 ± 0.518	-0.86 ± 0.353

No statistical analysis provided for Phase 2: Change in Body Weight at Week 36

Reported Adverse Events

No Adverse Events Entered.

More Information

Certain Agreements:

Principal Investigators are NOT employed by the organization sponsoring the study. There IS an agreement between Principal Investigators and the Sponsor (or its agents) that restricts the PTs rights to discuss or publish trial results after the trial is The agreement is: The only disclosure restriction on the PI is that the sponsor can review results communications prior to public release and can embarge communications retrial results for a period that is less than on equal to 60 days. The sponsor cannot require changes to the communication and cannot extend the embarge.

The only disclosure restriction on the PI is that the sponsor can review results communications prior to public release and can embargo communications regarding The cary discression of the Pris that the sported can review results communications price to point release and can emologic communications regards that results for a period that is more than 60 days but less than or equal to 180 days. The sponsor cannot require changes to the communication and cannot extend the embargo.

Other disclosure agreement that restricts the right of the PI to discuss or publish trial results after the trial is completed.

Limitations and Caveats

Limitations of the study, such as early termination leading to small numbers of participants analyzed and technical problems with measurement leading to unreliable or uninterpretable data

Results Point of Contact:

Name/Title: Chief Medical Officer Organization: Amylin Pharmaceuticals Inc e-mail: ctricalinals@amylin.com

No publications provided

Reportable Pairy.

Amplin Phermaceuticals (Lisa Proter, MD, Study Director)

Study Pina Received:

April 12, 2007

April 10, 2009

April 10, 2009

Clinical Trials gov Identifier.

Health Authority.

Responsible Party: Study ID Numbers: Study First Received: Results First Received:

